David Burstein

List of Publications by Year in descending order

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#	Article	IF	CITATIONS
1	Chimeric CRISPR-CasX enzymes and guide RNAs for improved genome editing activity. Molecular Cell, 2022, 82, 1199-1209.e6.	9.7	29
2	Engineered B cells expressing an anti-HIV antibody enable memory retention, isotype switching and clonal expansion. Nature Communications, 2020, 11, 5851.	12.8	42
3	A scoutRNA Is Required for Some Type V CRISPR-Cas Systems. Molecular Cell, 2020, 79, 416-424.e5.	9.7	49
4	Tiny Hidden Genes within Our Microbiome. Cell, 2019, 178, 1034-1035.	28.9	1
5	The distinction of CPR bacteria from other bacteria based on protein family content. Nature Communications, 2019, 10, 4173.	12.8	112
6	A Functional Mini-Integrase in a Two-Protein Type V-C CRISPR System. Molecular Cell, 2019, 73, 727-737.e3.	9.7	22
7	Hydrogen-based metabolism as an ancestral trait in lineages sibling to the Cyanobacteria. Nature Communications, 2019, 10, 463.	12.8	87
8	Mediterranean grassland soil C–N compound turnover is dependent on rainfall and depth, and is mediated by genomically divergent microorganisms. Nature Microbiology, 2019, 4, 1356-1367.	13.3	170
9	Programmed DNA destruction by miniature CRISPR-Cas14 enzymes. Science, 2018, 362, 839-842.	12.6	757
10	Genomic resolution of a cold subsurface aquifer community provides metabolic insights for novel microbes adapted to high CO ₂ concentrations. Environmental Microbiology, 2017, 19, 459-474.	3.8	212
11	Identical bacterial populations colonize premature infant gut, skin, and oral microbiomes and exhibit different in situ growth rates. Genome Research, 2017, 27, 601-612.	5.5	99
12	RNA Targeting by Functionally Orthogonal Type VI-A CRISPR-Cas Enzymes. Molecular Cell, 2017, 66, 373-383.e3.	9.7	229
13	Potential for microbial H2 and metal transformations associated with novel bacteria and archaea in deep terrestrial subsurface sediments. ISME Journal, 2017, 11, 1915-1929.	9.8	137
14	Retroelement-guided protein diversification abounds in vast lineages of Bacteria and Archaea. Nature Microbiology, 2017, 2, 17045.	13.3	62
15	New CRISPR–Cas systems from uncultivated microbes. Nature, 2017, 542, 237-241.	27.8	471
16	Novel Microbial Diversity and Functional Potential in the Marine Mammal Oral Microbiome. Current Biology, 2017, 27, 3752-3762.e6.	3.9	82
17	ldentification of novel <scp><i>X</i></scp> <i>anthomonas euvesicatoria</i> type <scp>III</scp> effector proteins by a machineâ€earning approach. Molecular Plant Pathology, 2016, 17, 398-411.	4.2	66
18	Two distinct RNase activities of CRISPR-C2c2 enable guide-RNA processing and RNA detection. Nature, 2016, 538, 270-273.	27.8	854

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19	Major bacterial lineages are essentially devoid of CRISPR-Cas viral defence systems. Nature Communications, 2016, 7, 10613.	12.8	224
20	Genomic analysis of 38 Legionella species identifies large and diverse effector repertoires. Nature Genetics, 2016, 48, 167-175.	21.4	235
21	Analysis of five complete genome sequences for members of the class Peribacteria in the recently recognized Peregrinibacteria bacterial phylum. PeerJ, 2016, 4, e1607.	2.0	57
22	Accurate, multi-kb reads resolve complex populations and detect rare microorganisms. Genome Research, 2015, 25, 534-543.	5.5	121
23	Novel Type III Effectors in Pseudomonas aeruginosa. MBio, 2015, 6, e00161.	4.1	37
24	Identification of Novel Coxiella burnetii Icm/Dot Effectors and Genetic Analysis of Their Involvement in Modulating a Mitogen-Activated Protein Kinase Pathway. Infection and Immunity, 2014, 82, 3740-3752.	2.2	55
25	Computational modeling and experimental validation of the <i>Legionella</i> and <i>Coxiella</i> virulence-related type-IVB secretion signal. Proceedings of the National Academy of Sciences of the United States of America, 2013, 110, E707-15.	7.1	166
26	DNA motifs determining the efficiency of adaptation into the <i>Escherichia coli</i> CRISPR array. Proceedings of the National Academy of Sciences of the United States of America, 2013, 110, 14396-14401.	7.1	62
27	CoPAP: Coevolution of Presence–Absence Patterns. Nucleic Acids Research, 2013, 41, W232-W237.	14.5	30
28	Changes in exon–intron structure during vertebrate evolution affect the splicing pattern of exons. Genome Research, 2012, 22, 35-50.	5.5	88
29	A Machine Learning Approach To Identify Hydrogenosomal Proteins in Trichomonas vaginalis. Eukaryotic Cell, 2012, 11, 217-228.	3.4	24
30	Uncovering the co-evolutionary network among prokaryotic genes. Bioinformatics, 2012, 28, i389-i394.	4.1	36
31	Differential GC Content between Exons and Introns Establishes Distinct Strategies of Splice-Site Recognition. Cell Reports, 2012, 1, 543-556.	6.4	249
32	Native homing endonucleases can target conserved genes in humans and in animal models. Nucleic Acids Research, 2011, 39, 6646-6659.	14.5	27
33	Genome-Scale Identification of Legionella pneumophila Effectors Using a Machine Learning Approach. PLoS Pathogens, 2009, 5, e1000508.	4.7	236
34	Large-scale comparative analysis of splicing signals and their corresponding splicing factors in eukaryotes. Genome Research, 2008, 18, 88-103.	5.5	161
35	The Average Common Substring Approach to Phylogenomic Reconstruction. Journal of Computational Biology, 2006, 13, 336-350.	1.6	184
36	Information Theoretic Approaches to Whole Genome Phylogenies. Lecture Notes in Computer Science, 2005, , 283-295.	1.3	6